

From the

INTERNATIONAL PRELIMINARY EXAMINING AUTHORITY

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NOTIFICATION OF TRANSMITTAL OF INTERNATIONAL PRELIMINARY **EXAMINATION REPORT**

(PCT Rule 71.1)

Date of Mailing

23 AUG 2004

(day/month/year) Applicant's or agent's file reference IMPORTANT NOTIFICATION ISPT-1000 International filing date (day/month/year) Priority date (day/month/year) International application No. 10 June 2002 (10.06.2002) PCT/US03/18003 09 June 2003 (09.06.2003) Applicant ISIS PHARMACEUTICALS, INC.

- The applicant is hereby notified that this International Preliminary Examining Authority transmits herewith the international preliminary examination report and its annexes, if any, established on the international application.
- 2. A copy of the report and its annexes, if any, is being transmitted to the International Bureau for communication to all the elected Offices.
- Where required by any of the elected Offices, the International Bureau will prepare an English translation of the report (but not of any annexes) and will transmit such translation to those Offices.

4. REMINDER

The applicant must enter the national phase before each elected Office by performing certain acts (filing translations and paying national fees) within 30 months from the priority date (or later in some Offices)(Article 39(1))(see also the reminder sent by the International Bureau with Form PCT/IB/301).

Where a translation of the international application must be furnished to an elected Office, that translation must contain a translation of any annexes to the international preliminary examination report. It is the applicant's responsibility to prepare and furnish such translation directly to each elected Office concerned.

For further details on the applicable time limits and requirements of the elected Offices, see Volume II of the PCT Applicant's Guide.

Name and mailing address of the IPEA/US

Mail Stop PCT, Attn: IPEA/US Commissioner for Patents P.O. Box 1450 Alexandria, Virginia 22313-1450

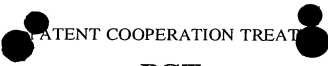
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Authorized officer

Telephone No. 571-272-1600

J. D. Schultz, Ph.D. A. Roberts for

Form PCT/IPEA/416 (July 1992)



PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

| Applicant's or agent's file reference | FOR FURTHER ACTION | | on of Transmittal of International examination Report (Form PCT/IPEA/416) |
|---|---|-------------------|--|
| ISPT-1000 International application No. | International filing date (day/n | | Priority date (day/month/year) |
| PCT/US03/18003 | 09 June 2003 (09.06.2003) | | |
| International Patent Classification (IPC) | | : | 10 0000 2002 (10.00.2002) |
| IPC(7): A61K 48/00; C07H 21/00; C120 | Q 1/68 and US Cl.: 514/44; 435 | /6, 325, 375; 536 | /23.1, 24.5 |
| Applicant | | | · |
| ISIS PHARMACEUTICALS, INC. | | | |
| Examining Authority and is | ary examination report has be stransmitted to the applicant a total of sheets, including | according to Ar | |
| which have been amer before this Authority (| nded and are the basis for this see Rule 70.16 and Section 6 | report and/or si | description, claims and/or drawings heets containing rectifications made nistrative Instructions under the PCT). |
| These annexes consist of a total of sheets. | | | |
| 3. This report contains indications relating to the following items: | | | |
| I Basis of the repor | rt | | |
| II Priority | | | |
| III Non-establishmen | III Non-establishment of report with regard to novelty, inventive step and industrial applicability | | |
| IV Lack of unity of i | nvention | | |
| V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement | | | |
| VI Certain document | | C | |
| VII Certain defects in the international application | | | |
| VIII Certain observations on the international application | | | |
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| Date of submission of the demand | Date | of completion o | f this report |
| 09 January 2004 (09.01.2004) | 08 Au | gust 2004 (08.08. | 2004) |
| Name and mailing address of the IPEA/US | Autho | rized officer | |
| Mail Stop PCT, Attn: IPEA/US Commissioner for Patents | J. D. | Schultz, Ph.D. | 7. Roberts for |
| P.O. Box 1450 Alexandria, Virginia 22313-1450 | | none No. 571-272 | |
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Form PCT/IPEA/409 (cover sheet)(July 1998)



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| Internationa | cation No. |
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| ٠. | : | | Internationa cation No. |
|----|----------|---|---|
| • | Ţ | NTERNATIONAL PREIS NARY EXAMINATION REPORT | PCT/US03/16663 |
| 4 | I. Ba | sis of the report | · · |
| | | th regard to the elements of the international application:* | |
| | | the international application as originally filed. | • |
| | | the description: | |
| | | pages 1-111 as originally filed | |
| | | pages NONE, filed with the demand pages NONE, filed with the letter of | · |
| | | the claims: | |
| | | pages 112 and 113 as originally filed | |
| | | pages NONE, as amended (together with any statem | ent) under Article 19 |
| | | pages NONE, filed with the demand pages NONE, filed with the letter of | |
| | | the drawings: | |
| | | pages None , as originally filed | |
| | | pages NONE, filed with the demand pages NONE, filed with the letter of | |
| | | _ | |
| | | the sequence listing part of the description: pages 1-35, as originally filed | · |
| | } | pages NONE filed with the demand | |
| | | pages NONE, filed with the letter of, filed with regard to the language, all the elements marked above were a | vailable or furnished to this Authority in the |
| | lan | guage in which the international application was filed, unless others elements were available or furnished to this Authority in the factors. | erwise indicated under this item. |
| | | the language of a translation furnished for the purposes of inter | |
| | | the language of publication of the international application (und | |
| | | the language of the translation furnished for the purposes of in | |
| | | 55.2 and/or 55.3). | |
| | 3. Wi | th regard to any nucleotide and/or amino acid sequence disclosemational preliminary examination was carried out on the basis of | sed in the international application, the fifthe sequence listing: |
| | | contained in the international application in printed form. | · |
| | | filed together with the international application in computer rea | adable form. |
| | | furnished subsequently to this Authority in written form. | |
| | | furnished subsequently to this Authority in computer readable | |
| | | The statement that the subsequently furnished written sequence international application as filed has been furnished. | |
| | | The statement that the information recorded in computer reada has been furnished. | ble form is identical to the written sequence listing |
| | 4. | The amendments have resulted in the cancellation of: | |
| | | the description, pages NONE | |
| | | | |
| | | the claims, Nos. NONE the drawings, sheets/fig NONE | |
| | | This report has been established as if (some of) the amendments had r | not been made, since they have been considered to go |
| | 5 | heyond the disclosure as filed, as indicated in the Supplemental Box (| Rule 70.2(c)).** |
| | thin was | acement sheets which have been furnished to the receiving Office in resport as "originally filed" and are not annexed to this report since they do replacement sheet containing such amendments must be referred to under | oonse to an invitation under Article 14 are referred to in o not contain amendments (Rules 70.16 and 70.17). |
| | ** Any | replacement sneet containing such unenanients must be rejerred to that | • |

cation No. Internationa PCT/US03/

| 1. STATEMENT | | |
|--------------------------------|-------------------|------|
| Novelty (N) | Claims 5-9, 15-20 | YES |
| 10.025 (1) | Claims 1-4, 10-14 | NO |
| Inventive Step (IS) | Claims 15-20 | YES |
| mitomitte step (15) | Claims 1-14 | NO |
| Industrial Applicability (IA) | Claims 1-21 | YES |
| industrial rippinousiary (1.2) | Claims NONE | NONO |

2. CITATIONS AND EXPLANATIONS

Claims 15-20 meet the criteria set out in PCT Article 33(2)-(3), because the prior art does not teach or fairly suggest methods of treatment using IRAK-1 targeted antisense compounds in methods of treating disease.

Claims 1-20 meet the criteria set out in PCT Article 33(4), and thus possess industrial applicability because the subject matter claimed can be made or used in industry.

Please See Continuation Sheet.

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V. 2. Citations and Explanations:

Claims 1-4, and 10-14 lack novelty under PCT Article 33(2) as being anticipated by Guo et al.

The invention of the above claims is drawn to modified antisense compounds 8 to 80 nucleobases long that hybridize with and inhibit IL-1 Receptor Associated Kinase-1 (IRAK-1), and methods of using same.

Guo et al. teaches modified antisense compounds 8 to 80 nucleobases long that hybridize with and inhibit IL-1 Receptor Associated Kinase-1 (IRAK-1), and methods of using same. See Materials and Methods of Guo.

Claims 1-14 lack an inventive step under PCT Article 33(3) as being obvious over the prior art as applied in the immediately preceding paragraph and further in view of Baracchini et al. and Taylor et al.

The invention of the above claims is drawn to antisense compounds that target IRAK-1 or said compounds comprising internucleoside, nucleobase, and 2' modifications, chimeras, or compositions comprising said compounds and pharmaceutically acceptable diluents thereof.

Guo et al teach the cDNA sequence encoding IRAK-1. Guo does not teach antisense sequences comprising nucleobase, and 2' modifications, and chimeras.

Taylor et al. teach that antisense oligonucleotides 7-30 nucleotides long can be synthesized to inhibit the expression of any protein provided the cDNA sequence is known. Taylor et al. also indicate that making and using such oligos are available to those of ordinary skill in the art, that it is common practice to chemically modify the such oligonucleotides to prolong their bioactivity, and also teach that with software analysis and high affinity oligos, one needs to screen only 3-6 oligos to find one that inhibits its target 66-95% (p. 565).

Baracchini et al. teach that antisense oligonucleotides can be used for research purposes, and also teach that preferred antisense oligonucleotides are modified in their sugar, backbone linkage and nucleobase composition (col. 6). Baracchini teaches that such modifications are desirable in antisense oligos because these modifications have desirable properties such as enhanced cellular uptake, enhanced affinity for nucleic acid targets and increased stability in the presence of nucleases. Baracchini et al provide specific embodiments of such modifications at columns 6-8 and in Example 1. These specific examples taught by Baracchini et al include the presently claimed phosphorothioate linkages, 2'-O-methoxyethyl sugars, 5-methylcytosine and chimeric oligonucleotides. Tables 1-4 show the successful design and use of modified oligonucleotides in cells in culture. Table 1 exemplifies the successful practice of general antisense design taught at columns 8-10. Column 4 teaches various carriers for antisense delivery. Baracchini et al. also teaches at column 8 that antisense oligonucleotides are preferably 8 to 30 nucleotides and that it is more preferable to make antisense oligonucleotides that are 12 to 25 nucleotides in length. Baracchini is considered to comprise a detailed blueprint for how to make and use inhibitory antisense oligos to target any known gene.

It would have been obvious to one of ordinary skill in the art to modify the antisense sequence of Guo to as taught by Taylor and Baracchini IRAK-1 expression for inhibition of IRAK-1 expression, and further, it would have been obvious to one of ordinary skill in the art to incorporate modifications as taught by Baracchini et al. a into said antisense compounds.

One would have been motivated to create such compounds because Guo et al. expressly teach antisense compounds that

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target and hybridize to IRAK-1. One would have been motivated to modify said antisense compounds as taught by Baracchini et al. a because they teach that such modifications increase an antisense compound's cellular uptake, target affinity and resistance to degradation.

Finally, one would have a reasonable expectation of success given that Taylor teaches that with software analysis and high affinity oligos, one needs to screen only 3-6 oligos to find one that inhibits its target 66-95%, and since Baracchini et al. and Bennett et al. both teach making modified antisense compounds targeted to distinct regions of a target gene, the steps of which are routine to one of ordinary skill in the art.

Thus in the absence of evidence to the contrary, the invention as a whole would have been prima facic obvious to one of ordinary skill in the art at the time the invention was made.

US 5,801,154 A (BARACCHINI et al.) 01 September 1998 (01.09.1998), entire document.

TAYLOR et al. Antisense oligonucleotides: a systematic high-throughput approach to target validation and gene function determination. Drug Disc. Today, 1999, Vol. 4, No. 12, pages 562-567, entire document.